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Authors

Hoffman, Adam S Fang, Chia-Yu Gates, Bruce C

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Homogeneity of Surface Sites in Supported Single-Site Metal Catalysts: Assessment with Band Widths of Metal Carbonyl Infrared Spectra

Adam S. Hoffman,^a; Chia-Yu Fang,^{a,b}; and Bruce C. Gates^a*

^{*a*}Department of Chemical Engineering, University of California at Davis, Davis, California, 95616, USA

^bDepartment of Materials Science and Engineering, University of California at Davis, Davis, California, 95616, USA

Corresponding Author

*Bruce C. Gates. Tel: (530) 752-3953 E-mail: bcgates@ucdavis.edu

ABSTRACT

Determining Identifying and controlling the uniformity of isolated metal sites on surfaces of supports are central goals in investigations of single-site catalysts, because well-defined species provide opportunities for fundamental understanding of the surface sites for fundamental understanding and industrial applications. CO is a useful probe molecule of the surface metal sites, often reacting with them to formleading to the formation of metal gem-dicarbonyls, the infrared spectra of which provide insights into the nature of the metal_sites and the metal_support interface. Metals bonded to various various support surface sites give broad bands in the spectra, and when narrow Full width at half maximum values characterizing the v_{co} values bands are observed, they provide evidence of indicate a highthe degree of uniformity of the metal sites, because metals bonded to a set of various sites give broad bands in the spectra. Much rReecent work focused on single-site catalysts has been done with uses supports that are inherently nonuniform, giving supported metal species that are therefore leading to a non-uniform metal sites, and ultimately resulting in a lack of fundamental understanding of the chemistry. Herein we summarize new and already reported values of v_{CO} data characterizing supported iridium *gem*dicarbonyls, showing that the most nearly uniform of them are those supported on zeolites and the least uniform are those supported on metal oxides. Guided by v_{CO} data of supported iridium *gem-dicarbonyls*, we have determined new, general synthesis methods to maximize the degree of uniformity of iridium speciesites on zeolites and on MgO. We that can be applied to other similar classes of support report results for a zeolite HY-supported iridium *gem*-dicarbonyl with full width at half maximum values of only 4.62 and 5.23 cm⁻¹ characterizing the symmetric and asymmetric CO stretches and implying that this is the most nearly uniform supported single-site metal catalyst.

TOC GRAPHIC



KEYWORDS-

Single-site catalyst, infrared spectroscopy, metal *gem*-dicarbonyl, surface heterogeneity, iridium, rhodium

Single-site supported catalysts typically incorporate metal cations bonded to support oxygen atoms, 1, -2, 3 and their catalytic properties depend markedly on the support. 4-9 and activities compared to the supported metal clusters Because almost all support surfaces are intrinsically heterogeneous, the supported species are nonuniform—bonded in various ways on various sites of the supports. Therefore, it is challenging to determine the structures of the supported species and the metal-support bondinginteraction bonding, and such information is incomplete in the above-mentioned work. The complexity of such these materials hinders fundamental understanding of their function as catalysts. This complexity also gives rise to varied catalytic activity of the materials. Physical characterization would provide the deepestmost understanding of the<u>sem_materials</u> if the supports were ideal—<u>that is</u>, perfectly crystalline materials with only single kinds of sites for bonding the metal. Herein we show how to assess the degree of uniformity of atomically dispersed metals on supports with infrared (IR) spectroscopy of CO ligands bonded to the metals, and guided by the physical characterization data, wewe report how to determined how to synthesize supported iridium complexes that are evidently appear to be the most nearly uniform of any single-site catalysts supported metal species. Thus, this work demonstrates improved synthesis methods guided by the results and understanding of physical measurements.

Values of v_{CO} characterizing a compound $M(CO)_xL_y$, where M is a transition metal and L a ligand, with the subscripts x and y taking on ranges of values, are sensitive to the backbonding involving *d*-electrons of the metal and π^* orbitals of the CO ligands. <u>TheFurther</u>, v_{CO} values characterize the electron-donor properties of L as well as the symmetry of the compound.

<u>Further, the v_{CO} values</u> <u>IR spectra similarly</u> provid<u>einge</u> essential details of the local environments of metal sites on solid surfaces.⁷¹³

The typical mononuclear metal carbonyl complex on a metal oxide support is <u>heterogeneous</u>, characterized by a smear of species and broad v_{CO} bands, but Miessner et al.^{8–14}_showed that mononuclear rhodium carbonyls bonded to dealuminated HY zeolite are characterized by narrow v_{CO} bands, indicating that the zeolite bonding sites are almost all the same. There are now numerous examples of single-site catalysts bonded to zeolites that broadly bear out Miessner's observations;;.^{3,159} <u>some are listed in Table 1.</u>

-Here we use full width at half maximum (FWHM) values of the carbonyl bands to classify supported metal complexes and show how this criterion <u>can be used to guidehelps to assess</u> the synthesis to <u>maximize the degree of uniformity of the of some of the most nearly uniform</u> supported metal species. We report new synthesis methods demonstrating how to take advantage of optimizing (a) synthesis temperature, (b) the zeolite support Si/Al ratio, and (c) the degree of crystallinity and surface site uniformity of a metal oxide support, MgO, <u>in order generate a higher, and in some cases, a nearly uniform supported Hr to maximize the degree of uniformity of supported iridium carbonyls.</u>-

Supported metal complexes were made from the precursor $Ir(CO)_2(acac)$ (acac is acetylacetonate, $C_5H_7O_2^{-}$) or $Ir(C_2H_4)_2(acac)$ and zeolite or metal oxide supports by reaction of the precursor with support surface OH groups to give supported $Ir(CO)_2^{16-2240-16}$ or $Ir(C_2H_4)_2$ groups; with the iridiummetal loadings of the samples being with metal loadings of -1.0 wt% <u>Hr</u>.^{16-19,2240-13,16} Supported $Ir(C_2H_4)_2$ groups weare readily converted to $Ir(CO)_2$ by reaction with CO. synthesis and characterization conditions were used in order to prevent metal site aggregation that occurs at higher temperatures.²³

In the reported room-temperature synthesis with zeolite HY,^{18,21+2+5} Ir(CO)₂(acac), which is barely soluble in *n*-pentane, evidently dissolv<u>esed</u> in this solvent, <u>was-is</u> transported to the zeolite, and react<u>sed</u> with zeolite OH groups. Syntheses with a wide range of supports have <u>similarly</u> given a family of supported Ir(CO)₂ complexes (Table 1), and a number of them have been characterized by extended X-ray absorption fine structure (EXAFS) data showing that each Ir atom (on average) bonds to 2 oxygen atoms of the support.^{18,21,22,2412,15-17} IR data characterizing the CO ligands confirm the presence of anchored iridium *gem*-dicarbonyls,^{18,19,21,22,2412,13,15-17} and DFT calculations²⁵¹⁰ support the structure assignments.-

___Now we report improved synthesis methods that give more nearly uniform supported species than those reported...

<u>includingWe report FWHM values of the carbonyl bands to demonstrate the uniformity</u>. Because the precursor Ir(CO)₂(acac) does not dissolve visibly in *n*-pentane solvent at room temperature, we carried out <u>a new synthesissome syntheses</u> at 193 K (the "low-temperature synthesis") in the following way: a flask containing *n*-pentane, zeolite HY particles, and Ir(CO)₂(acac) crystals was placed in a dry ice/isopropanol bath, and, over 24 h, the initially white zeolite particles turned light pink (Figure 1), and there was no observed change in the liquid color. Then the flask was slowly (over 30 min) warmed to room temperature, and the zeolite color turned to white/light gray (Figure 1). IR spectra consist of bands at 2109 and 2038 cm⁻¹ (Figure 2) characterizeing the symmetric and asymmetric stretching frequencies of terminal CO ligands, that—with the frequencies matching those previously–reported for single-site iridium carbonyls and; confirming the formation of a supported iridium *gem*-dicarbonyl. The lack of observed v_{CO} Failure to identify bands at lower-frequencies characteristic of bridging iridium carbonyls confirms the assignment to we are confident in the single-site assignmentspecies. -(The band characterizing the zeolite-supported sample and located between the iridium *gem*-dicarbonyl bands (Figure 2), at 2074 cm⁻¹, has been attributed to iridium tricarbonyl species, inferred, consistent with our weak bands, to be a minority species.²⁶) Need Reason for Tricarbonyl... cannot come up with one...-The band observed at 1870 cm⁻¹ observed in (Figure 2) for the Ir(CO)₂/HY zeolite is assigned to framework vibrations of the zeolite, and it is correspondingly present in the IR spectra of the support prior to addition the incorporation of the iridium.



Figure 1. Images characterizing samples at two stages of the low-temperature synthesis as $Ir(CO)_2(acac)$ reacted with zeolite HY to give a loading of 1.0 wt% Ir. Left: aAfter $Ir(CO)_2(acac)$ and zeolite HY had been slurried in *n*-pentane for 24 h in a dry ice bath at (193 K); right: after the slurry had been allowed to warm to 298 K in a period of approximately 30 min.

In a comparison experiment (the "<u>roomhigh</u>-temperature synthesis," matching that reported^{18,2112,15}), the reactants and solvent were slurried at room temperature, and the zeolite turned light gray in color as the Ir(CO)₂(acac) crystals dissolved and the precursor reacted with the zeolite. A comparison of the IR data characterizing the two samples (Table 1) shows that the frequencies of the CO bands characterizing the supported iridium *gem*-dicarbonyls were the same in both, with the symmetric stretch observed at 2109 cm⁻¹ and the asymmetric stretch at

2038 cm⁻¹, but the FWHM results differ, with the values for the low-temperature synthesis being 4.6 and 5.2 cm⁻¹ for v_{sym} and v_{asym} , respectively, and those for the <u>roomhigh</u>-temperature synthesis being 5.3 and 5.6 cm⁻¹, respectively. <u>Errors in the This comparison shows that the new low-temperature synthesis method gives a more nearly uniform sample than the roomhigh-temperature method. Deviation in the FWHM values offor the v_{sym} and v_{asym} were found to be \pm 0.2 and \pm 0.4 cm⁻¹, respectively, determined from data characterizing <u>based upon comparing</u> 7seven separately madeunique batches of Ir(CO)₂/zeolite HY. Thus, the comparison shows that the new low-temperature synthesis method gives more nearly uniform samples than the roomtemperature method. These errors show that the difference in the FWHM observed between the room- and low-temperature syntheses are unique and that synthesis temperature does influences the uniformity of the metal sites. ____(The band characterizing the zeolite-supported sample and located between the iridium *gem*-dicarbonyl bands (Figure 2), at 2074 cm⁻⁴, has been attributed to iridium tricarbonyl species.⁴⁹)</u>



Figure 2. IR spectra in the v_{CO} region of supported $Ir(CO)_2$ complexes demonstrating the variation FWHM as the uniformity of the bonding sites on the support surfaces decreases, roughly from top to bottom. $Ir(CO)_2$ on HY zeolite was made by the low-temperature method described in the text. The spectrum of $Ir(CO)_2/\gamma$ -Al₂O₃ is takenreproduced from the work of Lu et al.¹⁹

The results are consistent with the interpretation that the slow initial low-temperature adsorption of the precursor gave an initially nearly uniform distribution of iridium in the zeolite pores, followed by the higher-temperature reaction of the precursor with the zeolite surface to maintain such a distribution. We infer that in the low-temperature synthesis the precursor dissolved slowly in the *n*-pentane and diffused into the zeolite pores where it slowly became physisorbed before reacting, favoring an even distribution of metal in the zeolite and giving the pink sample with physisorbed precursor—and, when the temperature was later increased, these

species reacted with the zeolite surface OH groups, liberating acac ligands <u>16-2240-46</u> and turning the pink sample white with Ir(CO)₂ groups bonded to the zeolite. In contrast, we infer that in the <u>roomhigh</u>-temperature synthesis, the iridium complex reacted more quickly with the zeolite surface before it could spread evenly within the zeolite pores, thereby giving a less than uniform distribution of the iridium, which was likely concentrated near the pore mouths. <u>Further investigation with electron microscopy would be able to sort out the difference in metal distribution between the two synthesis temperatures, comparing the Ir bonded to the edges/amorphous regions[ref_Claudia_acs_catal_2014] and that in the pores, definitively determining if low temperature synthesis improves uniformity.</u>



Figure 1. Images characterizing samples at two stages of synthesis as $Ir(CO)_2(acac)$ reacted with zeolite HY to give a loading of 1.0 wt% Ir. Left: After $Ir(CO)_2(acac)$ and zeolite HY had been slurried in *n*-pentane for 24 h in a dry ice bath (193 K); right: after the slurry had been allowed to warm to 298 K in approximately 30 min.

For comparison of the supported species with dissolved molecular species, we determined FWHM values for the CO bands of the pure precursor Ir(CO)₂(acac) dissolved in mixed hexanes; the values are 3.9 and 4.0 cm⁻¹ for the symmetric and asymmetric stretches at 2074 and 2000 cm⁻¹, respectively (Table 1, Figure 2); as best we can tell, these data match within error those already reported for Ir(CO)₂(acac) in hexane.²⁰²⁷ The satellite-weak bands at 2056 and 1967 cm⁻¹ have been assigned to-as natural--abundance ¹³CO stretching frequenciesbands.²⁰²⁴

Most important for our work, the FWHM values of the dissolved iridium complex are only slightly less than those observed for the HY zeolite-supported sample made in the low-temperature synthesis (Figure 2)<u>compared to that in any other metal oxide</u>. The bands characterizing this zeolite-supported iridium *gem*-dicarbonyl are as narrow as any reported for such a supported species, and, to our knowledge, are consistent with the suggestion-inference that it is the most nearly uniform such-supported <u>single-site catalyst</u>sample.

Reports of numerous other supported metal carbonyl complexes are characterized by much broader bands than these (Table 1, Figure 2). For example, an γ-Al₂O₃-supported iridium complex is characterized by FWHM values that are an order of magnitude greater than those of the zeolite-supported sample (Table 1).



Figure 2. IR spectra in the v_{CO} region of supported $Ir(CO)_2$ complexes demonstrating the variation FWHM as the uniformity of the bonding sites on the support surfaces decrease, roughly from top to bottom. The $Ir(CO)_2/\gamma$ -Al₂O₃ is taken from the work of Lu et al.⁴³

To test another strategy for increasing the degree of uniformity of supported iridium carbonyls, we carried out a similar synthesis with a different zeolite, H-Beta, again at 298 K, with Ir(CO)₂(acac) as the precursor and the conditions stated above for the <u>roomhigh</u>-temperature synthesis. Two forms of zeolite H-Beta were used, one with a Si/Al atomic ratio of 19 and the otherone with a Si/Al atomic ratio of 150. The IR data give evidence of nearly the same supported Ir(CO)₂ groups in each of these zeolites, with the symmetric CO stretch observed at 2105 cm⁻¹ and the asymmetric stretch at 2033 cm⁻¹, but, significantly, the FWHM values were found to be 5.4 and 7.4 cm⁻¹, respectively, for the sample with the sparse population of Al sites and 10.1 and 14.9 cm⁻¹, respectively, for the other (Table 1). We suggest that the zeolite with the more widely dispersed sites has a more nearly uniform set of such sites (Al-OH groups) for bonding to the iridium and therefore gives a more uniform set of iridium-containing species, or, alternatively, that the chemisorption of the precursor on the sparsely distributed sites takes place more slowly than on the less sparsely distributed sites, giving a more uniform distribution of the iridium before chemisorption takes place. The band near 2070 cm⁻¹ corresponding to iridium tricarbonyl species was not observed in the zeolite beta samples. We speculate that the lack of band stems from zeolite beta having a higher degree of uniformity, whereas the pretreatment of zeolite Y generates amorphous regions adding non-uniformity to the sample.^{29ref} The lack of band near 2070 cm⁻⁺ indicated the presence of Ir(CO)₃ species compared to the species observed in the HY zeolite. This absences is likely due to size exclusion due to the pore size of H-Beta (6.68 Å) compared to HY (11.24Å) not allowing the formation of the tricarbonyl

We extended the synthesis method to a support with a much less uniform set of surface bonding sites than the zeolites, namely, MgO. The MgO samples were treated at various temperatures, 673, 1073, and 1273 K, under vacuum. These treatments lead to various degrees of bulk crystallinity, as shown by (a) differences in morphology observed by microscopy, ²²³⁰₋₋ (b) variations in the OH sites on the surface inferred from IR²³³¹₋₋ and electron paramagnetic resonance spectra, ²⁴³²₋₋ and (c) DFT²⁵³³₋₋ models showing various surface structures and surface sites. Our treatment procedures gave samples of MgO that differed from each other in terms of (1) the average crystallite domain size, as determined by X-ray diffraction crystallography (XRD) (Figure 3), and (2) the surface hydroxyl group density, as shown in the Supporting Information.



Figure 3. Correlation between the degree of crystallinity of the MgO support as measured by XRD and the degree of uniformity of the MgO-supported Ir(CO)₂ complexes measured by the FWHM of the symmetric CO stretching band determined by IR spectroscopy. Temperatures next to the data points represent the pretreatment temperature of the MgO support.

The samples of MgO-supported $Ir(CO)_2$, $Ir(CO)_2/MgO_T$, where *T* is the MgO pretreatment temperature in K, were synthesized at 298 K by the <u>roomhigh</u>-temperature method mentioned

above, except that the precursor was $Ir(C_2H_4)_2(acac)$. The synthesis gave supported $Ir(C_2H_4)_2$ species, as expected,¹⁶⁴⁰ and these were converted to $Ir(CO)_2$ by treatment with CO, as <u>expectedbefore</u>¹⁶⁴⁰ (Table 1). IR spectra show that the frequencies of the symmetric and asymmetric bands of the CO ligands depend on the MgO pretreatment temperature, *T*, consistent with the expectation that the surface bonding sites varied with *T*. Furthermore, the data demonstrate a decrease in the FWHM values of the v_{CO} bands with increasing *T* (Figure 2 and Table 1), as follows: for the symmetric band, the FWHM started at 23.9 cm⁻¹ and decreased to 22.5 cm⁻¹ and then to 16.4 cm⁻¹ as *T* increased from 673 to 1073 to 1273 K. A similar trend (FWHM values of 30.9, 30.4, and 15.5 cm⁻¹, respectively) was observed for the asymmetric stretch. Evidently the uniformity of the support bonding sites depends on the MgO pretreatment temperature, and the IR spectra demonstrate this point (although the structures of the bonding sites are still a matter of discussion).



Figure 3. Correlation between the degree of crystallinity of the MgO support as measured by the FWHM of the MgO (200) plane determined by XRD measurements and the degree of uniformity of the MgO-supported Ir(CO)₂ complexes measured by the FWHM of the symmetric

CO stretching band determined by IR spectroscopy. Temperatures next to the data points represent the pretreatment temperature of the MgO support.

The comparison shown in Table 1 of the FWHM values of the carbonyl bands in a family of supported Ir(CO)₂ complexes demonstrates a wide range of degrees of uniformity among the samples. Those having the highest degree in uniformity are zeolites, withand metal organic frameworks (MOFs) havingwith nodes that are small clusters of metal oxides arebeing less uniform than the zeolites, and bulk metal oxides beingare even less uniform, with those having a highly crystalline character, illustrated by MgO, being more nearly uniform than less crystalline solids illustrated by γ-Al₂O₃. Figure 3 illustrates the results for a representative family of these materials.

In summary, we have illustrated several synthetic approaches to control the degree of uniformity of single-site supported metal catalysts, by varying (1) the synthesis temperature, (2) the zeolite Si/Al ratio, and (3) the degree of metal oxide crystallinity guided by physical characterization. The comparison of the FWHM values characterizing CO ligands on the iridium inof a number of supported iridium carbonyls demonstrates a wide range of degrees of uniformity of the supported species and points to zeolites as the most nearly ideal reported supports for single-site catalysts with the low-temperature synthesis method giving most nearly uniform species reported—and these, we infer, provide some of the best opportunities for fundamental understanding of supported single-site metal catalysts...

Table 1. Comparison of supported single-site iridium and rhodium *gem*-dicarbonyl catalysts on the basis of the CO stretching frequencies and FWHM values, depending on the support and support treatment conditions, metal precursor, and synthesis conditions used to form the metal carbonyl.

| | Support | Support treatment | Precursor | Metal loading (wt %) | Synthesis conditions | ν _{co} (cm ⁻¹) | | Full width at half maximum of CO bands (cm ⁻¹) | | Other characterization | Reference |
|--|--|--|--|----------------------------|---|-------------------------------------|--------------------------|--|----------------------|--|-------------------------|
| | | | | | | Symmetric vibration | Asymmetric vibration | Symmetric vibration | Asymmetric vibration | techniques | |
| | None (Ir(CO)₂(acac) in- mixed hexane <u>s</u>) | | Ir(CO) ₂ (acac) | | | 2074 | 2000 | 3.9 | 4 | | This work |
| | H-Beta zeolite (Si/Al ~ 19) | O₂ 773 K 4 h; Vac. 773 K 16 h | Ir(CO) ₂ (acac) | 1 | Room temperature | 2105ª | 2033ª | 10.1 | 14.9 | | This work |
| | H-Beta zeolite (Si/Al ~ 150) | O ₂ 773 K 4 h; Vac. 773 K 16 h | Ir(CO)₂(acac) | 1 | Room temperature | 2105ª | 2033ª | 5.4 | 7.4 | | This work |
| | HY zeolite (Si/Al ~ 30) | O ₂ 773 K 2 h; Vac. 773 K 14 h | Ir(CO)₂(acac) | 1 | Room temperature | 2109 ^a | 2038ª | 5.3 | 5.6 | | This work |
| | HY zeolite (Si/Al ~ 30) | O ₂ 773 K 2 h; Vac. 773 K 14 h | Ir(CO) ₂ (acac) | 1 | <u>Low-</u> <u>temperatureCold</u> synthesis | 2109ª | 2038ª | 4.6 | 5.2 | | This work |
| | HY zeolite (Si/Al ~ 30) | O ₂ 773 K 2 h; Vac. 773 K 14 h | Ir(CO) ₂ (acac) | 1 | Room temperature | 2109ª | 2038ª | 5 | 5 | EXAFS spectroscopy; DFT calculation | <u>21</u> 15 |
| | H-SSZ-42 zeolite (Si/Al ~ 15) | O₂ 773 K 4 h; Vac. 773 K 16 h | Ir(C ₂ H ₄) ₂ (acac) | 1 | Room temperature; CO pulse | 2102ª; 2086 ^b | 2029ª; 2013 ^b | Broad | Broad | EXAFS spectroscopy | <u>19</u> 13 |
| | H-SSZ-53 zeolite | O ₂ 723 K 1 h; Vac. 723 K 1 h | Ir(C ₂ H ₄) ₂ (acac) | 1 | Room temperature; CO pulse | 2099ª | 2027ª | 10 | 10 | EXAFS spectroscopy | <u>18</u> 12 |
| | NaY zeolite (Si/Al ~ 2.6) | O ₂ 773 K 4 h; Vac. 773 K 12 h | Ir(CO)2(acac) | 1 | Room temperature | 2082 ^b | 1995 ^{<i>b</i>} | Broad | Broad | EXAFS spectroscopy | <u>24</u> 17 |

| | NU-1000 | Vac. 363 K 2h; Vac. 393 K 12h | Ir(CO) ₂ (acac) | 10 | Room temperature | 2066ª; 2082 ^b | 1990°; 2005 ^b | Broad | Broad | EXAFS spectroscopy; | <u>22</u> 16 |
|---|---|---|---|----|-------------------------------|---------------------------------------|---------------------------------------|-------|-------|--|------------------------|
| | | | | | | | | | | DFT calculation | |
| | NU-1000 | Vac. 363 K 2h; Vac. 393 K 12h | $Ir(C_2H_4)_2(acac)$ | 1 | Room temperature; CO pulse | 2066ª | 1900 ^a | Broad | Broad | EXAFS spectroscopy; DFT calculation | <u>22</u> 16 |
| | UiO-66 | Vac. 363 K 2h; Vac. 423 K 12h | Ir(CO)2(acac) | 10 | Room temperature | 2074 ^a ; 2085 ^b | 1996 ^a ; 2010 ^b | Broad | Broad | EXAFS spectroscopy; DFT calculation | <u>22</u> 16 |
| | UiO-66 | Vac. 363 K 2h; Vac. 423 K 12h | Ir(C ₂ H ₄) ₂ (acac) | 1 | Room temperature; CO pulse | 2074ª | 1996ª | Broad | Broad | EXAFS spectroscopy; DFT calculation | <u>2216</u> |
| | γ -Al ₂ O ₃ | O ₂ 773 K 2 h; Vac. 773 K overnight | $Ir(C_2H_4)_2(acac)$ | 1 | Room temperature; CO pulse | 2075ª | 1996 ^a | 30 | 30 | EXAFS spectroscopy | 13 17 |
| | MgO | Vac. 523 K 75 min; 673 K 60 min | Ir(C ₂ H ₄) ₂ (acac) | 1 | Room temperature; CO pulse | 2056ª | 1973ª | 23.9 | 30.9 | | This work |
| | MgO | Vac. 523 K 75 min; 1073 K 60 min | $Ir(C_2H_4)_2(acac)$ | 1 | Room temperature; CO pulse | 2061ª | 1983ª | 22.5 | 30.4 | | This work |
| | MgO | Vac. 523 K 75 min; 1273 K 60 min | $Ir(C_2H_4)_2(acac)$ | 1 | Room temperature; CO pulse | 2066ª | 1985ª | 16.4 | 15.5 | | This work |
| | None (Rh(CO) ₂ (acac) in <i>n</i> -hexane) | | Rh(CO) ₂ (acac) | | | 2084 | 2015 | 2.8 | 2.9 | | <u>27</u> 20 |
| | H-Beta zeolite (Si/Al ~ 19) | O₂ 773 K 16 h; Vac. 773 K 4 h | Rh(CO)2(acac) | 1 | Room temperature | 2115" | 2048ª | 9 | 9 | EXAFS spectroscopy | 26<u>34</u> |
| 1 | H-SSZ-42 zeolite (Si/Al ~ 15) | O ₂ 723 K 1 h; Vac. 723 K 1 h | Rh(CO) ₂ (acac) | 1 | Room temperature | 2111ª; 2082 ^b | 2045ª; 2020 ^b | Broad | Broad | EXAFS spectroscopy | <u>34</u> 26 |
| | H-Mordenit <u>e</u> e zeolite | O₂ 773 K 16 h; Vac. 773 K 4 h | Rh(CO)₂(acac) | 1 | Room temperature | 2111ª; 2092 ^b | 2045ª; 2035 ^b | Broad | Broad | EXAFS spectroscopy | <u>34</u> 26 |
| | HY zeolite (Si/Al ~ 30) | O ₂ 773 K 4 h; Vac. 773 K 16 h | Rh(C ₂ H ₄) ₂ (acac) | 1 | Room temperature; CO pulse | 2117ª | 2052ª | < 8 | < 8 | EXAFS spectroscopy; DFT calculation | <u>35</u> 27 |
| z | Ultra-stable Y eolite (Si/Al ~ 95) | | [Rh(NH ₃) ₅ Cl](OH) ₂ | c | d | 2118ª | 2053ª | < 5 | < 5 | - | <u>14</u> 8 |

^{*a*} Assigned as chemisorbed species.

^{*b*} Assigned as physisorbed species.

^c The metal loading was not specified.

^{*d*} The synthesis was carried out by an ion-exchange method. Ultra-stable Y zeolite was mixed with $[Rh(NH_3)_5Cl](OH)_2$ in an aqueous solution. The sample was dried at 383 K for 3 h and calcined in air at 673 K for 2 h. The calcined sample then reacted with CO (at 10 Torr and 423 K) for 30 min, followed by subsequent evacuation at 300 K.

EXPERIMENTAL METHODS

Syntheses. Room-temperature synthesis of Ir(CO)₂ on zeolite HY, H-Beta, or MgO followed was done with methods reported elsewhere.^{16-19,21,22,24,34,35} ^{10-13,15-17,26,27} The supports were pretreated at temperatures from 773 to 1273 K in-an O20xygen and/or under vacuum atmosphere-to remove excess moisture and carbon--containing compounds. The precursor, Ir(CO)₂(acac) (Sigma-Aldrich) or $Ir(C_2H_4)_2(acac)$ (synthesized in this work²⁸³⁶), and support (, HY zeolite (Zeolyst CBV 760), H-Beta zeolite (Zeolyst, CP814C and CP814C-300) or MgO (EMD)) were slurried in npentane (Sigma-Aldrich, 98%) then evacuated to remove the solvent. Low-temperature synthesis of Ir(CO)₂ on zeolite HY was achieved by placing a Schlenk flask containing the metal precursor, $Ir(CO)_2(acac)$, zeolite powder, and *n*-pentane in a Dewar filled with a bath containing dry ice and isopropanol, with the slurry stirred for 24 h. The flask was then removed and allowed to warm to room temperature before the *n*-pentane was removed by evacuation. The air- and moisturesensitive samples were stored in an argon-atmosphere glovebox. In an argon- atmosphere, tThe supported $Ir(C_2H_4)_2$ samples were packed in a flow reactor or flow-through cell, transferred to a flow system without exposure to air, and then-and exposed to CO (Airgas 10% in helium) to form the Ir(CO)₂ species. The carbonyl species were recovered in the glovebox.

IR spectroscopy. A Bruker IFS-66VS spectrometer was used to collect spectra of samples that had been loaded into an air-tight <u>flow-through</u> cell in an argon-atmosphere glovebox to minimize oxygen and moisture exposure. <u>CO exchange was conducted as spectra were being recordedim</u>-<u>situ to monitor the conversion from the iridium diethylene to the iridium dicarbonyl complex</u>. Each spectrum is an average of 64 scans collected over 2 min. <u>Details of the experiments with</u> solid samples are reported elsewhere.^{16-19,21,22,24,34,35} The spectrum of Ir(CO)₂(acac) in hexanes (EM Science, 98.5%) was measured with a saturated solution in a sealed liquid IR cell (International Crystal Laboratories) with a 0.5--mm path length.

Associated Content

Supporting Information contains the IR spectra and XRD diffraction patterns of the MgO supports pretreated at various temperatures <u>and EXAFS data characterizing a zeolite-supported</u> <u>iridium gem-dicarbonyl</u>.

Corresponding Author

Corresponding Author: Bruce C. Gates. Tel: (530) 752-3953 E-mail: bcgates@ucdavis.edu

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